

SPECIAL HEALTH SERVICES MEDICAL CONDITIONS Revised: 5-22-2019

ACQUIRED BRAIN INJURY

ACUTE FLACCID MYELITIS (AFM)

ADENOID HYPERTROPHY causing SLEEP APNEA

ALPHA 1-ANTITRYPSIN DEFICIENCY

AMPUTATION

AMYOTONIA CONGENITA requiring rehabilitative measures

ANAL STENOSIS & IMPERFORATE ANUS

ANEMIAS (excluding minor anemias), including sickle cell

APLASIA CUTIS CONGENITA, severe, requiring surgery &

ECTODERMAL DYSPLASIA

ARNOLD-CHIARI DEFORMITY

ARTHROGRYPOSIS

ASTHMA, persistent, requiring controller medications

ATAXIAS, FAMILIAL DEGENERATIVE DISEASE requiring

rehabilitative measures

ATTENTION-DEFICIT/HYPERACTIVITY DISORDER

(ADD/ADHD)

AUTO-IMMUNE DISORDERS, chronic, severe, and complex in

nature

BILE DUCT ATRESIA

BIRTH INJURY (ERB's PALSY, etc.) requiring bracing or

surgery

BONE CYST requiring surgery

BONE TUMORS, benign, requiring surgery, including

OSTEOCHONDROMAS

BONY DEFORMITIES requiring bracing, casting or surgery & POST-TRAUMATIC DEFORMITY (orthopedic or severe

soft tissue deformity due to injury; excluding acute fracture

without an underlying condition)

BRAIN TUMORS requiring surgery and/or radiation

BRANCHIOGENIC CLEFT CYST requiring surgery

BREAST HYPOPLASIA causing considerable psychological

problems requiring surgery

BURNS, severe, acute, including residuals

CANCER, including CANCER OF EYE

CATARACTS

CELIAC DISEASE

CEREBRAL PALSY, congenital or acquired, requiring

rehabilitative measures

CHOANAL ATRESIA

CHRONIC LUNG/LOWER AIRWAY CONDITIONS, including

chronic lung disease, chemical pneumonitis, and subglottic

stenosis

CLEFT LIP AND/OR PALATE, including SHORT PALATE

and SUBMUCOUS CLEFT

COLLAGEN VASCULAR DISORDERS, including but not

limited to lupus, dermatomyositis, scleroderma, Sjogren's

syndrome, and rheumatoid arthritis

CONGENITAL ADRENAL HYPERPLASIA (CAH)

CORNEAL TRANSPLANTS

CRANIOSTENOSIS (premature synostosis)

CYSTIC FIBROSIS

CYSTIC HYGROMA

CYSTINOSIS

DENTAL DISORDERS, congenital

DERMATOMYOSITIS

DIABETES INSIPIDUS

DIABETES MELLITUS, TYPE I and TYPE II

DIAPHRAGMATIC HERNIA

DISLOCATION OF HIPS OR OTHER JOINTS

EAR DEFORMITY

EHLERS-DANLOS DISEASE

ENCEPHALITIS, POLIOMYELITIS OR MENINGITIS,

residuals of

ENUCLEATION (removal of eyeball)

EOSINOPHILIC GASTROENTERITIS

EPIDERMOLYSIS BULLOSA

ESOPHAGEAL VARICES

EYE WOUNDS, penetrating

EYELID DEFORMITY requiring surgery, congenital

FACE DEFORMITY

FEMORAL CAPITAL EPIPHYSIS, slipped

GASTROINTESTINAL TRACT ANOMALIES, congenital

(including gastroschisis)

GENITO-URINARY TRACT ANOMALIES, congenital,

severe and requiring surgery

GENU RECURVATUM, severe

GENU VALGUM (Knock-knees), severe

GENU VARUM (Bowed legs), severe

GLAUCOMA, congenital

GROWTH HORMONE DEFICIENCY

GUILLAIN-BARRE DISEASE, severe, acute, requiring

tracheotomy and/or ventilation, including residuals

HEARING LOSS

HEART CONDITIONS, congenital or acquired

HEMANGIOMA, medically significant

HEMOGLOBINOPATHIES, limited to:

Sickle cell anemia

Thalassemia

HEMOPHILIA including deformities

HISTIOCYTOSIS X (eosinophilic granuloma)

HYDROCEPHALUS requiring surgery

HYPERCHOLESTEROLEMIA, congenital, including familial

combined hyperlipidemia

HYPERTHYROIDISM

HYPOPARATHYROIDISM, congenital or if suspected to last

longer than two years

HYPOPHOSPHATEMIC RICKETS

HYPOPITUITARISM

HYPOTHALAMIC ADRENAL INSUFFICIENCY

HYPOTHYROIDISM

ICHTHYOSIFORM ERYTHRODERMA, congenital, severe

IMMUNODEFICIENCY STATES including severe combined

immunodeficiency (SCID)

INFLAMMATORY BOWEL DISEASE including Crohn's

Disease and ulcerative colitis

INTERSEX DISORDERS, congenital

JOINT DEFORMITY, CLUBFEET AND CLUBHANDS,

severe, requiring bracing, casting, surgery or physical

therapy

JUVENILE IDIOPATHIC ARTHRITIS, (Juvenile 3-hydroxy-3-methylglutaryl-CoA lyase deficiency Rheumatoid Arthritis) Glutaric acidemia/aciduria Isobutyryl-CoA dehydrogenase deficiency Isovaleric acidemia (IVA) KYPHOSIS, adolescent, requiring bracing or surgery Methylmalonic acidemia (MMA) LARYNGEAL PAPILLOMA Propionic Acidemia Mitochondrial acetoacetyl-CoA thiolase deficiency LEUKEMIA LEUKODYSTROPHY, including adrenoleukodystrophy (BKT, 3-ketothiolase deficiency) Multiple CoA carboxylase deficiency Refsum's Disease (Phytanic acid restriction) MALOCCLUSION, handicapping MASTOIDITIS, chronic **MICROCEPHALY** MEGACOLON requiring surgery MUCOPOLYSACCHARIDOSIS (MPS) (including variants) METABOLIC DISORDERS/INBORN ERRORS OF **METABOLISM** NARCOLEPSY (with or without Cataplexy) Amino Acid Disorders, limited to: NEPHROSIS & CHRONIC NEPHRITIS Arginase deficiency/Hyperargininaemia NERVE INJURIES, chronic Argininemia **NEUROFIBROMATOSIS** Argininosuccinic acidemia (ASA lyase deficiency) NEUROMUSCULAR DISORDERS limited to those covered Carbamoyl phosphate synthetase deficiency by MDA including muscular dystrophy Citrullinemia (ASA synthetase deficiency) NEVI with malignant potential Glutaric acidemia/aciduria Glutathione synthetase deficiency (5-oxoprolinuria) OCULAR ALBINISM, congenital Homocystinuria (cystathione synthase deficiency) OSTEOCHONDRITIS of various bones Hypermethioninemia OSTEOGENESIS IMPERFECTA Hyperornithinemia, hyperammonemia, OSTEOMYELITIS, residuals of Homocitrullinemia (HHH syndrome) Hyperornithinemia or ornithine oxo-acid PANTOTHENATE KINASE-ASSOCIATED aminotransferase deficiency NEURODEGENERATION (PKAN) Maple syrup urine disease (MSUD) (Hallervorden-Spatz Disease, including infusion pump) N-Acetylglutamate synthetase deficiency PARAPLEGIA, traumatic, and its direct complications Nonketotic hyperglycinemia PECTUS CARINATUM/PECTUS EXCAVATUM requiring Ornithine aminotransferase deficiency surgery Ornithine transcarbamylase deficiency (OTC) PERTHES DISEASE Phenylketonuria (PKU), includes phenylalanine POLYCYSTIC KIDNEY DISEASE hydroxylase deficiency (PAH) and PRECOCIOUS PUBERTY hyperphenylalaninemia **PSEUDOHYPOPARATHYROIDISM** Tyrosinemia (I, II, III); **PSORIASIS** Biotinidase Deficiency PTOSIS (drooping eyelids) Fatty Acid Oxidation Disorders, limited to: PULMONARY LOBAR EMPHYSEMA 2,4 dienoyl-CoA reductase deficiency Long chain acyl-CoA dehydrogenase deficiency RETINAL DETACHMENT in Marfan's syndrome (LCADD) RETROLENTAL FIBROPLASIA (retinopathy of prematurity) Long chain 3-OH acyl-CoA dehydrogenase deficiency (LCHAD) SCLERODERMA Carnitine/acylcarnitine translocase deficiency (CACT) SCOLIOSIS requiring bracing or surgery Carnitine palmitoyltransferase deficiency-type I SEIZURE DISORDERS, excluding febrile seizures (CPTI) SHORT BOWEL SYNDROME Carnitine palmitoyltransferase deficiency-type II SPINA BIFIDA, MENINGOCELE, MYELOCELE STRABISMUS through age 10 (CPTII) Carnitine transport defect (CTD) SUBLUXATED EYE LENS in Marfan's syndrome Glutaric acidemia/aciduria SUPERNUMERARY PARTS, severe Medium chain acyl-CoA dehydrogenase deficiency SYNDACTYLY SYNDROMES, limited, requiring ongoing medical treatment; (MCAD) Multiple acyl-CoA dehydrogenase deficiency includes septo-optic dysplasia (MADD) or glutaric acidemia-type II (GAII) Short chain acyl-CoA dehydrogenase deficiency THROMBOCYTOPENIA, congenital (SCAD) (ethylmalonic academia) THROMBOEMBOLISM Trifunctional protein deficiency (TFP Deficiency) THYROGLOSSAL DUCT CYST T-LYMPHOCYTE IMMUNE DEFICIENCY STATE Very long chain acyl-CoA dehydrogenase deficiency (VLCAD) TORTICOLLIS (not spasmodic, requiring casting or surgery) Galactosemia TRACHEAL STENOSIS GLUT 1 Deficiency (glucose 1 transporter deficiency) TRACHEOESOPHAGEAL FISTULA

GLUT I Denciency (glucose I transporter denciency)
Glycogen Storage Disease
Hereditary Fructose Intolerance
Organic Acid Disorders, limited to:
2-methylbuyryl-CoA dehydrogenase deficiency
3-methylcrotonyl-CoA carboxylase deficiency
3-methylglutaconic-CoA hydratase deficiency

TRACHEAL STENOSIS
TRACHEOESOPHAGEAL FISTULA
TRANSVERSE MYELITIS
TUBERCULOSIS OF BONES AND JOINTS
TUBEROUS SCLEROSIS
UNDESCENDED TESTES
VASCULAR ABNORMALITIES
WEGENER'S GRANULOMATOSIS